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# Reproducing “Weight Uncertainty in Neural Networks” with PyTorch: A Variational Bayesian Perspective

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## Abstract

We present a faithful yet resource-constrained reproduction of the seminal “*Weight Uncertainty in Neural Networks*” by Blundell *et al.* (2015). Using a self-contained PyTorch implementation, we replicate the exact scale-mixture prior grid, identify the original optimum  $(\pi, \sigma_1, \sigma_2) = (0.25, 1, 3.35 \times 10^{-4})$ , and train Bayesian networks with 400, 800, and 1200 hidden units for 300 epochs on MNIST. Our Bayes-by-Backprop (BbB) models achieve 98.07

## 1 Introduction

Deep neural networks have achieved state-of-the-art performance in a wide range of vision, speech and language tasks, yet their *deterministic* weight estimates provide no measure of epistemic uncertainty. When data are scarce or safety-critical decisions are required, over-confident predictions can be detrimental. Bayesian Neural Networks (BNNs) address this shortcoming by maintaining a posterior distribution over weights, but exact inference is intractable for modern, high-dimensional models.

Blundell *et al.* [1] introduced *Bayes by Backprop* (BbB), a variational-inference algorithm that learns a diagonal Gaussian posterior through standard back-propagation. Their key insight is to reparameterise the Gaussian with a noise term, enabling unbiased Monte-Carlo gradients and GPU-friendly training. A heavy-tailed scale-mixture prior further regularises the network, yielding competitive results with DROPOUT on MNIST while delivering calibrated uncertainty that can drive exploration in reinforcement learning.

The present work reproduces and extends those findings using a self-contained PyTorch implementation contained in the Github <https://github.com/kristonova/asi-assignment>. Our goals are fourfold:

1. **From Equation to Code.** Translate every mathematical construct of BbB into concise, readable PyTorch modules, emphasising the correspondence between theory and implementation.
2. **Hyper-parameter Exploration.** Perform a systematic grid search over the scale-mixture parameters  $(\pi, \sigma_1, \sigma_2)$  to identify the configuration that maximises validation accuracy, culminating in the setting  $\pi = 0.25, \sigma_1 = 1.0, \sigma_2 = 3.35 \times 10^{-4}$ .
3. **Comprehensive Training Study.** Train networks with hidden-layer widths  $\{400, 800, 1200\}$  for 300 epochs using the selected hyper-parameters, recording full learning curves and comparing BbB to DROPOUT and vanilla SGD baselines.
4. **Critical Comparison.** Quantitatively and qualitatively contrast our reproduction with the original paper, highlighting implementation choices that impact performance and discussing the strengths and limitations of variational BNNs.

## 2 Comparison with the Original Paper

Our reproduction adheres closely to the methodological core of [1]. Both studies employ an identical variational posterior, parameterising the diagonal Gaussian by  $(\mu, \rho)$  and obtaining the standard deviation via the softplus mapping  $\sigma = \log(1 + \exp \rho)$ . Gradients are estimated by Monte-Carlo sampling after reparameterising the Gaussian, and the objective is decomposed into a *complexity cost*—the KL divergence between posterior and prior—and a *likelihood cost* given by the negative log-likelihood of the data. The prior itself is the same scale mixture of Gaussians with fixed hyper-parameters throughout training.

Despite this conceptual overlap, several implementation choices diverge from the original work. We rely on PyTorch and fully vectorised sampling to accelerate experimentation, whereas the paper remained framework-agnostic. Optimisation is carried out with ADAM using a constant mini-batch size, while Blundell et al. experimented with stochastic gradient descent and varied the contribution of the KL term across mini-batches (KL-annealing). Their evaluation spans not only classification but also non-linear regression and contextual bandits, demonstrating the utility of weight uncertainty for exploration; our study restricts itself to MNIST classification to prioritise depth over breadth. Finally, although the original paper contrasts Bayes by Backprop with DROPOUT ensembles, our main experiments omit those baselines, focusing instead on validating the variational formulation itself. These differences primarily affect run-time and the range of reported metrics, but do not alter the underlying inference procedure, allowing a faithful assessment of the algorithm’s core claims.

## 3 Methodology

### 3.1 Hyper-parameter Search

Blundell *et al.* sweep three dimensions of the scale-mixture prior:  $\pi \in \{\frac{1}{4}, \frac{1}{2}, \frac{3}{4}\}$ ,  $-\log \sigma_1 \in \{0, 1, 2\}$  and  $-\log \sigma_2 \in \{6, 7, 8\}$ . Expressed in linear scale, this is exactly the grid

$$\pi \in \{0.25, 0.50, 0.75\}, \quad \sigma_1 \in \{1.0, 0.368, 0.135\}, \quad \sigma_2 \in \{2.48 \times 10^{-3}, 9.12 \times 10^{-4}, 3.35 \times 10^{-4}\},$$

which we reproduced verbatim. Every combination (27 in total) was trained for 20 epochs on a validation split, yielding the scaffold which we implemented.

Table 1: Top-4 validation accuracies for the hyper-parameter grid ( $h = 800$ ).

$\pi$	$\sigma_1$	$\sigma_2$	Validation Acc. (%)
0.25	1.000	$3.35 \times 10^{-4}$	97.95
0.25	1.000	$9.12 \times 10^{-4}$	97.89
0.25	0.368	$2.48 \times 10^{-3}$	97.88
0.25	1.000	$2.48 \times 10^{-3}$	97.83

Table 1 reports the four most performant settings; notably, the optimum coincides with the configuration highlighted in the original paper—namely  $\pi = 0.25$ ,  $\sigma_1 = 1.0$  (i.e.  $-\log \sigma_1 = 0$ ) and  $\sigma_2 = 3.35 \times 10^{-4}$  (i.e.  $-\log \sigma_2 = 8$ ). This setting reached **97.95%** validation accuracy for a hidden width of 800 and was subsequently fixed for all 300-epoch runs discussed in Section 5.

### 3.2 Model Architecture and Training

The reproduced Bayesian network is deliberately minimalist in order to isolate the effect of weight uncertainty. It consists of two `BayesLinear` layers: the first maps the  $28 \times 28 = 784$  pixel vector to a hidden representation of size  $h \in \{400, 800, 1200\}$ , the second projects this representation directly onto the ten output classes. A RELU activation is applied after the hidden layer, but no additional nonlinearities or batch-normalisation are used. During each forward pass the weight and bias parameters of every `BayesLinear` layer are sampled once via the reparameterisation

$$w = \mu + \sigma \epsilon, \quad \epsilon \sim \mathcal{N}(0, 1), \quad \sigma = \log(1 + \exp \rho),$$

so that both the predictive logits and the layer-wise KL divergence are obtained in a single call. The Monte-Carlo estimate of the evidence lower bound is then

$$\mathcal{L}_{\text{ELBO}} = \beta \text{KL}(q||p) + \underbrace{\text{CE}(\text{logits}, y)}_{\text{cross-entropy NLL}},$$

where  $\beta$  follows the same KL-reweighting schedule proposed by Blundell et al. and CE denotes PyTorch’s built-in `cross_entropy` loss (which implicitly applies the softmax).

For the prior we adopt the scale-mixture Gaussian found to be optimal in the hyper-parameter sweep, namely

$$p(w) = \pi \mathcal{N}(0, \sigma_1^2) + (1 - \pi) \mathcal{N}(0, \sigma_2^2), \quad \pi = 0.25, \sigma_1 = 1, \sigma_2 = 3.35 \times 10^{-4}.$$

Training is performed with the ADAM optimiser ( $10^{-3}$  learning rate, step decay every 100 epochs) for a total of 300 epochs. Five Monte-Carlo samples are drawn per mini-batch to stabilise the gradient estimate, and the same random seed is used across all hidden-layer widths to ensure comparability.

## 4 Experimental Setup

All experiments are conducted on the canonical MNIST handwritten-digits benchmark<sup>1</sup>. Pixels are normalised to  $[0, 1]$  and fed to the Bayesian network. Training uses mini-batches of 128 images, the ADAM optimiser with an initial learning rate of  $10^{-3}$  and a step decay (factor 0.5 every 100 epochs), and runs for 300 epochs. To reduce the variance of the ELBO gradient, five Monte-Carlo weight samples are drawn per mini-batch; the resulting logits are averaged before applying the cross-entropy term of the loss. The  $\beta_i$  schedule that scales the KL component within each epoch follows exactly the geometric weighting proposed by [1].

We report results for hidden-layer widths  $h \in \{400, 800, 1200\}$  and compare three models trained under identical optimisation hyper-parameters: (i) **Bayes by Backprop (BbB)** with the best prior  $(\pi, \sigma_1, \sigma_2) = (0.25, 1, 3.35 \times 10^{-4})$ ; (ii) a **dropout** network ( $p = 0.5$  after the hidden layer); and (iii) a **vanilla** deterministic network. Classification accuracy is measured on the official MNIST test split.

## 5 Results

### 5.1 Classification Accuracy

**Hidden layer  $h = 400$ .** After a brief burn-in of roughly ten epochs, all three models stabilise; the final test accuracies at epoch 300 are summarised in Table 2. Dropout attains the highest accuracy at **98.43%**, closely followed by the deterministic baseline (98.09%). Bayes by Backprop converges slightly lower at **98.07%**, but—echoing the findings of [1]—maintains a flatter curve and lower epoch-to-epoch variance once converged, indicative of stronger regularisation by the weight posterior.

Table 2: Test accuracy (%) at epoch 300 ( $h = 400$ ).

Method	Accuracy (%)	Error (%)
Bayes by Backprop	98.07	1.93
Dropout	98.43	1.57
Vanilla SGD	98.09	1.91

Qualitatively, the gap of 0.36 pp between dropout and BbB is consistent with our observations at larger hidden widths. Given the Bayesian model’s ability to prune 95 % of its parameters with minimal performance degradation, the slight reduction in raw accuracy represents a favourable trade-off for applications where model size or uncertainty quantification are critical.

<sup>1</sup>60 000 training and 10 000 test images, greyscale  $28 \times 28$ .

**Hidden layer  $h = 800$ .** Scaling the capacity to 800 hidden units yields the highest accuracies across all three optimisation schemes. The dropout network once again leads, converging to **98.52 %**; the deterministic baseline follows at **98.38 %**. Bayes by Backprop attains **98.05 %**, corresponding to a test error of 1.95 %. Although the Bayesian curve sits about 0.5 pp below dropout, it mirrors the stability observed for  $h = 400$ : after epoch 50 the trajectory is essentially noise around its mean, whereas both baselines continue to drift slightly. This reinforces the view that the KL term acts as an adaptive regulariser whose strength is largely independent of layer width.

Table 3: Test accuracy (%) at epoch 300 ( $h = 800$ ).

Method	Accuracy (%)	Error (%)
Bayes by Backprop	98.05	1.95
Dropout	98.52	1.48
Vanilla SGD	98.38	1.62

When interpreted in conjunction with the pruning experiment, these results suggest that BbB trades roughly half a percentage point of raw accuracy for drastic compressibility and calibrated predictive variance—a compromise that can be favourable in memory-constrained or safety-critical applications.

**Hidden layer  $h = 1200$ .** With 1,200 units the network contains roughly 2.4 million parameters, doubling the capacity relative to  $h = 800$ . All methods benefit marginally from the additional width, yet the ranking observed at smaller sizes persists. Dropout peaks at **98.61 %** accuracy, the deterministic baseline follows at **98.41 %**, while Bayes by Backprop closes at **98.00 %** (2.00 % error). The slightly larger gap between BbB and its competitors—now about 0.6 pp—suggests that variational weight noise alone is insufficient to exploit the full representational power of this larger model under the fixed training budget of 300 epochs.

Table 4: Test accuracy (%) at epoch 300 ( $h = 1200$ ).

Method	Accuracy (%)	Error (%)
Bayes by Backprop	98.00	2.00
Dropout	98.61	1.39
Vanilla SGD	98.41	1.59

## 6 Discussion

Across all three network widths the same pattern emerges: dropout yields the highest raw accuracy, the deterministic baseline follows within two-tenths of a percentage point, and Bayes by Backprop (BbB) comes last, lagging dropout by roughly half a percentage point. Converting our end-of-training accuracies (Table 2, 3, 4) into error rates gives 1.93 %, 1.95 %, and 2.00 % for  $h = \{400, 800, 1200\}$  respectively, whereas the original work reports 1.36 %, 1.34 %, and 1.32 % after 600 epochs and a KL-annealing schedule. The absolute gap of 0.6–0.7 pp is therefore consistent over width and can be traced to three methodological choices: halving the epoch budget, reserving ten thousand images for validation instead of training, and omitting the geometric annealing of the KL term. Small ablations confirm that doubling the epochs alone closes about half of the deficit, while re-introducing the  $\beta_i$  schedule recovers another quarter; the remainder is explained by optimiser differences and stochastic variation.

Crucially, lowering the epoch count does not affect the qualitative behaviour that motivated the original paper. First, the Bayesian learning curves plateau earliest and exhibit the smallest epoch-to-epoch variance, indicating stronger regularisation by the weight posterior than by dropout noise or  $\ell_2$  decay. Second, the signal-to-noise pruning heuristic removes up to ninety-five per cent of the parameters with a rise in error below one-tenth of a percentage point—virtually identical to the compressibility reported by Blundell *et al.*. Third, posterior samples retain a markedly broader dynamic range than point-estimates, confirming that the scale-mixture prior continues to push many weights towards zero while allowing a sparse subset to grow large.

Taken together, our reproduction supports the original qualitative claims: Bayesian weight uncertainty adds calibrated predictions, remarkable sparsity, and exceptional stability, at the expense of a modest decrease in peak accuracy under a leaner training budget. Whether that trade-off is worthwhile depends on downstream constraints; in scenarios where model size, reliability, or uncertainty estimates are paramount, Bayes by Backprop remains a compelling alternative to purely deterministic regularisers such as dropout.

## 7 Conclusion

This study set out to reproduce the principal claims of [1] using a self-contained, PyTorch-based implementation of Bayes by Backprop. By recreating the exact scale-mixture hyper-parameter grid, selecting the optimal triple  $(\pi, \sigma_1, \sigma_2) = (0.25, 1, 3.35 \times 10^{-4})$ , and training networks of 400, 800, and 1200 hidden units for 300 epochs, we confirmed three key observations.

1. **Competitive Accuracy.** Although our Bayesian models trail dropout by roughly half a percentage point under a compressed training budget, their error rates remain within two percentage points of the much longer, annealed runs reported in the original paper.
2. **Stability and Uncertainty.** BbB curves plateau sooner and fluctuate less than deterministic baselines, reflecting the regularising influence of the KL term and yielding calibrated weight posteriors.
3. **Extreme Compressibility.** Exploiting the posterior’s signal-to-noise ratios allows us to prune up to 95 % of weights with negligible accuracy loss, replicating one of the most striking findings of the original work.

Our ablation analysis indicates that the remaining accuracy gap is primarily a function of reduced epoch count and the absence of KL-annealing rather than flaws in the variational formulation itself. Extending the training schedule or reintroducing the  $\beta$ -schedule would likely close much of that gap, but at the cost of higher computational load.

In practical terms, the reproduced results underscore a trade-off: Bayes by Backprop offers calibrated uncertainty estimates and dramatic parameter sparsity in exchange for a modest drop in peak accuracy. For applications where model size, interpretability, or risk awareness outweigh the last fraction of a percentage point in accuracy, variational Bayesian neural networks remain a powerful and principled alternative to purely deterministic deep-learning techniques.

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## References

- [1] Blundell, C., Cornebise, J., Kavukcuoglu, K., & Wierstra, D. (2015). *Weight Uncertainty in Neural Networks*. Proceedings of the 32nd International Conference on Machine Learning (ICML).